TLE 'HOME' ENTERED AT 22:36:37 ON 22 AUG 2008 => file caplus biosis COST IN U.S. DOLLARS SINCE FILE TOTAL ENTRY SESSION FULL ESTIMATED COST 0.21 0.21 FILE 'CAPLUS' ENTERED AT 22:36:58 ON 22 AUG 2008 USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT. PLEASE SEE "HELP USAGETERMS" FOR DETAILS. COPYRIGHT (C) 2008 AMERICAN CHEMICAL SOCIETY (ACS) FILE 'BIOSIS' ENTERED AT 22:36:58 ON 22 AUG 2008 Copyright (c) 2008 The Thomson Corporation => "positive charged (s) antigen" L1 0 "POSITIVE CHARGED (S) ANTIGEN" => positive (w) charged 21095 POSITIVE (W) CHARGED => antigen L3 902001 ANTIGEN => L2 (1) L3 281 L2 (L) L3 => L2 (p) L3 281 L2 (P) L3 => L2 (s) L3 73 L2 (S) L3 L6 => HCV L7 42824 HCV => L7 (s) L6 1 L7 (S) L6 => L7 (L) L6 1 L7 (L) L6 L9 => L6 and L7 L10 1 L6 AND L7 => adiuvant L11 110859 ADJUVANT => L11 and L6 16 L11 AND L6 L12 => D L8 IBIB ABS L8 ANSWER 1 OF 1 CAPLUS COPYRIGHT 2008 ACS on STN ACCESSION NUMBER: 2001:396697 CAPLUS DOCUMENT NUMBER: 135:4467 TITLE: Vaccine compositions INVENTOR(S): Drane, Debbie; Cox, John; Houghton, Michael; Paliard, Xavier

Csl Limited, Australia; Chiron Corporation

PATENT ASSIGNEE(S):

SOURCE: PCT Int. Appl., 67 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

LANGUAGE: Englis FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

	PA:	FENT	NO.			KIND DATE				APPI	LICAT	ION I	DATE					
						A1 20010531 A9 20020718					WO 2	2000-		20001117				
		W: AE, AG, AL,																
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	EP	1239	876			B1		2008	0730									
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			IE,	SI,	LT,	LV,	FI,	RO,	MK,	CY,	AL,	TR						
		5189				A							20001117					
	JP	2003	5148	72		Т		2003	0422		JP 2	2001-		20001117				
		5209								NZ 2000-520976								
										ZA 2002-3986 US 2003-622470							0020	
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PRIOR	RIORITY APPLN. INFO.:											1999-						
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											WO 2	2000-2	AU 1 4	10		W 2	0001	117

AB The present invention relates generally to an immunogenic complex comprising a charged organic carrier and a charged antigen and, more particularly, a neg. charged organic carrier and a pos. charged antigen, wherein the charged antigen is a polyprotein of Hepatitis C Virus (HCV), particularly the core protein of HCV, or a fragment thereof, or a fusion protein comprising the polyprotein or a fragment thereof. The complexes of the present invention are useful in vaccine compns. as therapeutic and/or prophylactic agents for facilitating the induction of immune responses, and in particular a cytotoxic T-lymphocyte response, in the treatment of a disease condition which results from an HCV infection.

REFERENCE COUNT: 9 THEER ARE 9 CITED REFERENCES AVAILABLE FOR THIS

REFERENCE COUNT: 9 THERE ARE 9 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

## => D L12 IBIB ABS 1-16

L12 ANSWER 1 OF 16 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 2004:975071 CAPLUS

DOCUMENT NUMBER: 142:416911

TITLE: Structure and adsorption properties of commercial

calcium phosphate adjuvant

AUTHOR(S): Jiang, Dongping; Premachandra, Gnanasiri S.; Johnston,

Cliff; Hem, Stanley L.

CORPORATE SOURCE: Department of Industrial and Physical Pharmacy, Purdue

University, West Lafayette, IN, 47907-2091, USA

SOURCE: Vaccine (2004), 23(5), 693-698 CODEN: VACCDE; ISSN: 0264-410X

PUBLISHER: Elsevier B.V. Journal DOCUMENT TYPE:

LANGUAGE: English

Calcium phosphate adjuvant is a com. available vaccine

adjuvant that potentiates the immune response to antigens.

Although its name suggests that it is Ca3(PO4)2, x-ray diffraction, FTIR spectroscopy, thermal anal, and the Ca/P molar ratio identify com, calcium phosphate adjuvant as non-stoichiometric hydroxyapatite, Cal0-x

(HPO4)x (PO4)6-x (OH)2-x, where x varies from 0 to 2. The surface charge is pH-dependent (point of zero charge = 5.5). Consequently, com. calcium phosphate adjuvant exhibits a neg. surface charge at physiol. pH

and electrostatically adsorbs pos. charged

antigens. The presence of hydroxyls allows calcium phosphate

adjuvant to adsorb phosphorylated antigens by ligand exchange with surface hydroxyls. 17 REFERENCE COUNT: THERE ARE 17 CITED REFERENCES AVAILABLE FOR THIS

L12 ANSWER 2 OF 16 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 2002:350514 CAPLUS

DOCUMENT NUMBER: 138:95296

TITLE: Adjuvant effect of zinc on microencapsulated tetanus toxoid

AUTHOR(S):

CORPORATE SOURCE:

McHugh, C.; Somavarapu, S.; Atuah, K.; Eyles, J.; Alpar, O.

Centre for Drug Delivery Research, University of

London School of Pharmacy, Bloomsbury, London, WC1N 1AX, UK

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

SOURCE: Proceedings - 28th International Symposium on Controlled Release of Bioactive Materials and 4th

Consumer & Diversified Products Conference, San Diego, CA, United States, June 23-27, 2001 (2001), Volume 2, 1083-1084. Controlled Release Society: Minneapolis,

Minn. CODEN: 69CNY8

DOCUMENT TYPE: Conference

LANGUAGE: English

The object of this study was to investigate the adjuvant effect of zinc (as zinc oxide) on the adjuvanticity of microencapsulated tetanus toxoid. We also investigated the effect of surface modification of the zinc oxide particles in order to increase the immune response obtained

using zinc alone as an adjuvant. The microencapsulation of zinc oxide with the antigen was found to increase the immune response

generated and coating the zinc oxide particles in order to make them more pos, charged was found to increase the immune titer

compared with uncoated particles.

REFERENCE COUNT: THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS 3 RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L12 ANSWER 3 OF 16 CAPLUS COPYRIGHT 2008 ACS on STN

2002:314786 CAPLUS ACCESSION NUMBER:

DOCUMENT NUMBER: 136:324057

TITLE: Vaccine composition comprising an adjuvant

/carrier peptide which enhances immune response to a

co-administered antigen

INVENTOR(S): Fritz, Joerg; Mattner, Frank; Zauner, Wolfgang; Nagy,

Eszter; Buschle, Michael

PATENT ASSIGNEE(S): Cistem Biotechnologies G.m.b.H., Austria SOURCE: PCT Int. Appl., 40 pp. CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1 PATENT INFORMATION:

	TENT I				KIND DATE					APPLICATION NO.							DATE			
									WO 2001-EP12041											
	W: AE, AG, AL,				AM.	AT.	AU.	AZ.	BA.	BE	3,	BG.	BR.	BZ.						
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711	2002	1122	26		2		2002	0423		7II	20	102-	1220	490	20011010					
717	2002	2122	26		D2		2002	0105		no	20	102-	1232	20001018 20011018 20011018						
ED	1226	212J.	20		3.1		2000	0716	EP 2001-980496						20011018					
EP	1326634				D1		2003	0/10	EP 2001-980496								0011	010		
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EC	2263	260			73 20060929					FC 2001-900490						20011010				
DII	2328	305			13		2000	0710	DII 2001-300430						20011010					
77	2003	0014	5.5		7		2000	0224	73 2003 1465						20030334					
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MA	2003	711100	000		A 20050304					IN 2003-MN263						20030226				
NO	2003	0016	020		A 20030/14					MX 2003-PA2828						20030331				
110	2005	0013	270		7.1		2005	0000		NO 2003-1595						2	0030	417		
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AB The invention relates to a vaccine which comprises at least one antigen and a peptide comprising a sequence R1-XXXANXXX+R2, whereby N is a whole number between 3 and 7, preferably 5, -X is a pos. charged natural and/or non-natural amino acid residue, Z is an amino acid residue selected from the group consisting of L, V, I, F and/or W, and Rl and R2 are selected independently from the other from the group consisting of -H, -NH2, -COCH3, -COH, a peptide with up to 20 amino acid residues or a peptide reactive group or a peptide linker with or without a peptide; X-R2 may also be an amide, ester or thioester of the C-terminal amino acid residue, as well as the use of said peptide for the preparation of an adjuvant for enhancing the immune response to at least one antigen. One example discusses the efficiency of peptide KIKLILLIKLKI in delivery of influenza hemaglutinin antigen to antigen-presenting cells and uptake into APCs. The peptide also enhances T-cell responses (i.e. interferon-y production) to the

antigen.

REFERENCE COUNT: 3 THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L12 ANSWER 4 OF 16 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 2001:396697 CAPLUS

DOCUMENT NUMBER: 135:4467

TITLE: Vaccine compositions

INVENTOR(S): Drane, Debbie; Cox, John; Houghton, Michael; Paliard,

PATENT ASSIGNEE(S): Csl Limited, Australia; Chiron Corporation

SOURCE: PCT Int. Appl., 67 pp.

CODEN: PIXXD2
DOCUMENT TYPE: Patent

LANGUAGE: English
FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

					KIND DATE								DATE				
WO	2001	A1 20010531 A9 20020718			1												
		CR, HU, LU, SD, ZA, GH,	CU, ID, LV, SE, ZW GM,	CZ, IL, MA, SG,	DE, IN, MD, SI,	DK, IS, MG, SK,	DM, JP, MK, SL,	DZ, KE, MN, TJ,	EE, KG, MW, TM,	ES KP MX TR	, BG, , FI, , KR, , MZ, , TT, , TZ, , LU,	GB, KZ, NO, TZ, UG,	GD, LC, NZ, UA,	GE, LK, PL, UG,	GH, LR, PT, UZ,	GM, LS, RO, VN,	HR, LT, RU, YU,
AU AU EP	7726	BJ, 843 0137: 17 876	CF,	CG,	CI, A1 A B2 A1	GA, 2001 2001 2004 2002	GN, 0531 0604 0506 0918	GW, ML, MR, NE, SN, CA 2000-2391843 AU 2001-13730 EP 2000-975681						20001117 20001117			
NZ ZA	5189 2003 5209 2002 2004	IE, 99 5148 76 0039:	SI, 72 86 270	LT,	LV, A T A	FI,	RO, 2002 2003 2005 2003	MK, 1220 0422 0128 1217	CY,	AL NZ JP NZ ZA US US US US	, IT, , TR 2000- 2001- 2002- 2002- 2003- 1999- 2000- 2000- 2000-	5189 5394 5209 3986 6224 1666 2243 7144	99 83 76 70 52P 62P 38	1 1 1	2 2 2 2 2 2 2 P 1 P 2 B1 2	0001 0001 0001 0020 0030 9991	117 117 117 520 721 119 811

more particularly, a neg. charged organic carrier and a pos. charged antigen, wherein the charged antigen is a polyprotein of Hepatitis C Virus (HCV), particularly the core protein of HCV, or a fragment thereof, or a fusion protein comprising the polyprotein or a fragment thereof. The complexes of the present invention are useful in vaccine compns. as therapeutic and/or prophylactic agents

AB The present invention relates generally to an immunogenic complex comprising a charged organic carrier and a charged antigen and,

for facilitating the induction of immune responses, and in particular a cytotoxic T-lymphocyte response, in the treatment of a disease condition which results from an HCV infection.

REFERENCE COUNT: 9 THERE ARE 9 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L12 ANSWER 5 OF 16 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 2000:786836 CAPLUS

DOCUMENT NUMBER: 134:339301

TITLE: Evaluation of a liposome-supplemented intranasal influenza subunit vaccine in a murine model system:

Induction of systemic and local mucosal immunity de Haan, Aalzen; van Scharrenburg, Guus J. M.; Masihi, AUTHOR(S):

K. Noel; Wilschut, Jan

CORPORATE SOURCE: Department of Medical Microbiology, Molecular Virology Section, University of Groningen, Groningen, 9713 AV,

SOURCE: Journal of Liposome Research (2000), 10(2 & 3),

159-177

CODEN: JLREE7; ISSN: 0898-2104

PUBLISHER: Marcel Dekker, Inc.

DOCUMENT TYPE: Journal LANGUAGE: English

The study reports on the mucosal immunoadjuvant activity of liposomes in an exptl. influenza subunit vaccine administered intranasally (i.n.) to mice. Antibody responses induced by the i.n. liposomal vaccine were compared to those induced by an influenza infection or by s.c. injection of subunit antigen alone, the conventional route of human flu vaccination.

Neg. charged liposomes, but not pos. charged or zwitterionic liposomes, co-administered i.n. with influenza subunit

antigen, stimulated systemic IgG levels and local antibody responses in pulmonary secretions, relative to the responses upon i.n. administration of subunit antigen alone. I.n. immunization with liposome-supplemented subunit antigen as well as s.c. immunization with subunit antigen alone or infection induced high levels of IgG antibodies in serum and pulmonary secretions, with a preferential induction of IgG1

upon immunization and IgG2a upon infection. Both i.n. immunization with liposome-supplemented antigen and infection, but not s.c. immunization with subunit antigen alone, induced local secretion of S-IgA. At the same time, both IgA- and IgG-secreting cells appeared in the lungs and

lung-associated lymph nodes, suggestive of local antibody production Thus, the liposomal adjuvant system, combined with a mucosal

administration protocol, provides a promising strategy for induction of both systemic and local antibody responses against influenza virus. REFERENCE COUNT: 48 THERE ARE 48 CITED REFERENCES AVAILABLE FOR THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L12 ANSWER 6 OF 16 CAPLUS COPYRIGHT 2008 ACS on STN ACCESSION NUMBER: 2000:592580 CAPLUS

DOCUMENT NUMBER: 133:191986

TITLE: Immunogenic complexes and methods relating thereto

INVENTOR(S): Cox, John Cooper; Drane, Debbie Pauline PATENT ASSIGNEE(S): CSL Limited, Australia

SOURCE: PCT Int. Appl., 111 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent. LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT I	KIN	D	DATE			APPL	ICAT	DATE								
WO 2000	A1 200008			0824		WO 2		20000217								
W:	ΑE,	AL,	AM,	AT,	AU,	AZ,	BA,	BB,	BG,	BR,	BY,	CA,	CH,	CN,	CR,	CU,
	CZ,	DE,	DK,	DM,	EE,	ES,	FI,	GB,	GD,	GE,	GH,	GM,	HR,	HU,	ID,	IL,
	IN,	IS,	JP,	KE,	KG,	KP,	KR,	KZ,	LC,	LK,	LR,	LS,	LT,	LU,	LV,	MA,
	MD,	MG,	MK,	MN,	MW,	MX,	NO,	NZ,	PL,	PT,	RO,	RU,	SD,	SE,	SG,	SI,

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SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZW
         RW: GH, GM, KE, LS, MW, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF,
             CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
                         A1 20000824 CA 2000-2362204
     CA 2362204
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     AU 2000026515
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    AU 783344
                        B2 20051020
A1 20011107
     EP 1150710
                                          EP 2000-904734
         R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
             IE, SI, LT, LV, FI, RO
     JP 2002537271 T 20021105
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    NA 513935
ZA 2001006521
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                              20030310
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PRIORITY APPLN. INFO.:
                                           AU 1999-8735
                                                              A 19990217
                                           AU 1999-1861
                                                              A 19990727
                                           WO 2000-AU110
                                                              W 20000217
    The present invention relates generally to an immunogenic complex
     comprising a charged organic carrier and a charged antigen and,
     more particularly, a neg. charged organic carrier and a pos.
    charged antigen. The complexes of the present invention
     are useful, inter alia , as therapeutic and/or prophylactic agents for
     facilitating the induction of a cytotoxic T-lymphocyte response to an
     antigen.
REFERENCE COUNT:
                              THERE ARE 5 CITED REFERENCES AVAILABLE FOR THIS
                              RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT
L12 ANSWER 7 OF 16 CAPLUS COPYRIGHT 2008 ACS on STN
ACCESSION NUMBER:
                       2000:526177 CAPLUS
DOCUMENT NUMBER:
                        134:250883
TITLE:
                        Influence of antigenic forms and adjuvants
                        on protection against a lethal infection of Aujeszky's
                        disease virus
                       Katayama, S.; Oda, K.; Ohqitani, T.
AUTHOR(S):
CORPORATE SOURCE:
                      Division of Veterinary Microbiology, Kyoto Biken
                       Laboratories, Uji, Kyoto, 611-0041, Japan
SOURCE:
                        Vaccine (2000), 19(1), 54-58
                        CODEN: VACCDE; ISSN: 0264-410X
PUBLISHER:
                        Elsevier Science Ltd.
DOCUMENT TYPE:
                        Journal
LANGUAGE:
                        English
    The influence of antigenic forms and adjuvant types on
     protection against a lethal infection of Aujeszky's disease virus (ADV) in
    mice was investigated. Antiviral IgG2a antibody response against
     particulate (inactivated ADV) and soluble antigen (ADV solubilized
     with deoxycholate-Na) in approx. order of extent was ISA70 > QS-21 >
     pos. charged liposome > neg. charged liposome > weak
    neg. charged liposome > ISA25 > lablabside F saponin > aluminum phosphate
     gel > non adjuvant. Particulate antigen induced higher IgG2a
     antibody production than soluble antigen. Particulate antigen combined
     with ISA70, ISA25 or pos. charged liposome gave 100,
     50 and 40% protection to mice, resp. In contrast, soluble antigen plus ISA70
    conferred 30% protection on mice. Immunogens using the other
     adjuvants gave ≤20% protection to mice. These results
    indicate that a combination of particulate antigen and an appropriate
     adjuvant effectively induces the production of antiviral IgG2a
     antibody and provides protection against a lethal ADV infection in mice.
REFERENCE COUNT:
                        35
                              THERE ARE 35 CITED REFERENCES AVAILABLE FOR THIS
                              RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT
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L12 ANSWER 8 OF 16 CAPLUS COPYRIGHT 2008 ACS on STN ACCESSION NUMBER: 1999:557808 CAPLUS

AB

DOCUMENT NUMBER: 131:341812

TITLE: Positively charged liposome functions as an efficient

immunoadjuvant in inducing cell-mediated immune

response to soluble proteins
AUTHOR(S): Nakanishi, T.; Kunisawa, J.; Hayashi, A.; Tsutsumi,

Y.; Kubo, K.; Nakaqawa, S.; Nakanishi, M.; Tanaka, K.;

Mayumi, T.

CORPORATE SOURCE: Graduate School of Pharmaceutical Sciences, Osaka

University, Osaka, Japan

SOURCE: Journal of Controlled Release (1999), 61(1-2), 233-240

CODEN: JCREEC; ISSN: 0168-3659
PUBLISHER: Elsevier Science Ireland Ltd.

DOCUMENT TYPE: Journal

LANGUAGE: English

AB In order to design an optimized liposome immunoadjuvant for inducing cell-mediated immune response against soluble proteinaceous antigens, we investigated the effect of liposomal surface charge on the immunoadjuvant action. Pos. charged liposomes containing soluble

antigens functioned as a more potent inducer of antigen -specific cytotoxic T lymphocyte responses and delayed type

hypersensitivity response than neg. charged and neutral liposomes containing the same concus. of antigens. To clarify the reason of the differential immune response, we examined the delivery of soluble proteins by

the liposomes into the cytoplasm of macrophages, using fragment A of diphtheria toxin (DTA) as a marker. We found that pos. charged liposomes encapsulating DTA are cytotoxic to macrophages, while empty pos. charged liposomes, DTA in neg. charged and neutral liposomes are not. Consistent

with this, only macrophages pulsed with OVA in pos. charged liposomes could significantly stimulate OVA-specific, class I MHC-restricted T cell hybridoma. These results suggest that the pos. charged

liposomes can deliver proteinaceous antigens efficiently into the cytoplasm of the macrophages/antigen-presenting cells, where the antigens are processed to be presented by class I MHC mols.

to induce the cell-mediated immune response. Possible development of a safe and effective vaccine is discussed.

REFERENCE COUNT: 38 THERE ARE 38 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L12 ANSWER 9 OF 16 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 1998:497103 CAPLUS DOCUMENT NUMBER: 129:274369

ORIGINAL REFERENCE NO.: 129:55933a,55936a

TITLE: Cationization of liposomal surface charge enhances adjuvant effect of liposomes for tumor vaccine

AUTHOR(S): Nakanishi, Tsuyoshi; Kunisawa, Jun; Hayashi, Akira;
Tsutsumi, Yasuo; Hayakawa, Takao; Mayumi, Tadanori
CORPORATE SOURCE: Graduate School of Pharmaceutical Science, Osaka

University, Suita, Osaka, 565-0871, Japan

SOURCE: Yakuzaigaku (1998), 58(2), 59-68 CODEN: YAKUA2; ISSN: 0372-7629

PUBLISHER: Nippon Yakuzai Gakkai

DOCUMENT TYPE: Journal LANGUAGE: English

AB In order to design an optimum liposome immunoadjuvant for tumor vaccines, we investigated the relationship between liposome surface charge and adjuvant action. Pos. charged multilamellar vesicles (MLVs) were

taken up efficiently by macrophages, while neg. charged and neutral MLVs

were hardly picked up. Consistent with this, pos.

charged MLVs containing soluble ovalbumin (OVA) functioned as a more potent inducer of antigen-specific cytotoxic T lymphocyte (CTL)

responses and antibody production than neg. charged and neutral MLVs containing

the same concns. of antigens. Furthermore, the in vivo anti-tumor effects of variously charged liposomal antigens were examined using a Meth A tumor model and a crude butanol extract derived from Meth A (Meth A-CBE) as the tumor-associated antigen. Mice vaccinated with pos. charged MLVs containing Meth A-CBE showed significant inhibition of Meth A tumor growth compared to mice vaccinated with Meth A-CBE alone or mice vaccinated with neutral or neg. charged liposomal Meth A-CBE. The injection of carrageenan into mice led to a significant loss of anti-tumor vaccinal effect of pos. charged liposomal Meth A-CBE, which may be due to the inhibition of uptake and antigen

presentation of liposomal antigens by macrophages as a result of a lack of macrophages in the immune site. Our results indicate that the

pos. charge on the surface of liposomes represents an important factor for enhancing their immunoadjuvancy in the induction of antigen-specific immune responses and vaccinal effects against tumors.

REFERENCE COUNT: 23 THERE ARE 23 CITED REFERENCES AVAILABLE FOR THIS

RECORD, ALL CITATIONS AVAILABLE IN THE RE FORMAT

L12 ANSWER 10 OF 16 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 1997:775527 CAPLUS

DOCUMENT NUMBER: 128:66384

ORIGINAL REFERENCE NO.: 128:12895a,12898a

TITLE: Positively charged liposome functions as an efficient immunoadjuvant in inducing immune responses to soluble

proteins AUTHOR(S):

Nakanishi, Tsuyoshi; Kunisawa, Jun; Hayashi, Akira; Tsutsumi, Yasuo; Kubo, Kazuyoshi; Nakagawa, Shinkasu; Fujiwara, Hiromi; Hamaoka, Toshiyuki; Mayumi, Tadanori

CORPORATE SOURCE: Faculty and Graduate School of Pharmaceutical Science, Osaka University, Suita, 565, Japan

SOURCE: Biochemical and Biophysical Research Communications

(1997), 240(3), 793-797

CODEN: BBRCA9; ISSN: 0006-291X

PUBLISHER: Academic Press

DOCUMENT TYPE: Journal LANGUAGE: English

To design an optimum liposome immunoadjuvant for soluble protein antigens, the authors investigated the relationship between liposomal surface charge and adjuvant action. Pos. charged multilamellar vesicles (MLV) were taken up efficiently by macrophages, while neg. charged and neutral MLVs were hardly picked up. Consistent with this, pos. charged MLVs containing soluble chicken egg albumin (OVA) functioned as a

more potent inducer of antigen-specific cytotoxic T lymphocyte (CTL) responses and antibody production than neg. charged and neutral MLVs containing the same concns. of antigens. The pos. charge on the

surface of liposomes represents an important factor for enhancing their immunoadjuvancy in the induction of antigen-specific immune responses.

REFERENCE COUNT: 25 THERE ARE 25 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L12 ANSWER 11 OF 16 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 1997:351527 CAPLUS DOCUMENT NUMBER: 127:85895

ORIGINAL REFERENCE NO.: 127:16405a,16408a TITLE: Adjuvanticity and protective immunity elicited by Leishmania donovani antigens encapsulated in

positively charged liposomes AUTHOR(S):

Afrin, Farhat; Ali, Nahid CORPORATE SOURCE: Leishmania Group, Indian Institute Chemical Biology,

Calcutta, 700032, India

SOURCE: Infection and Immunity (1997), 65(6), 2371-2377 CODEN: INFIBR; ISSN: 0019-9567 American Society for Microbiology

DOCUMENT TYPE: Journal LANGUAGE: English

PUBLISHER:

AR In the search for a leishmaniasis vaccine, extensive studies of cutaneous leishmaniasis have been carried out. Investigations in this regard with the visceral form are limited. As an initial step in the identification of the protective mols., leishmanial antigens extracted from the membranes of

Leishmania donovani promastigotes, alone or in association with liposomes, were evaluated for their immunogenicity and ability to elicit a protective immune response against challenge infection. I.p. immunization of hamsters and BALB/c mice with the leishmanial antigens conferred protection against infection with the virulent promastigotes.

Encapsulation in pos. in pos. charged liposomes

significantly enhanced the protective efficacy of these antigens The splenic parasite burden of hamsters was reduced by 97% after 6 mo of infection. BALB/c mice exhibited 87 and 81.3% protection in the liver and spleen, resp., after 4 mo of infection. These protected animals elicited profound delayed-type hypersensitivity and increased levels of Leishmania-specific IqG antibodies. Protection in mice also coincided with elevated levels of IqM and IqA antibodies, which decreased with disease progression in the control-infected animals. Although both IGG1 and IgG2a antibodies were present in the sera of infected mice, IgG1 appeared to be the predominant isotype, suggesting a preferential

induction of the Th2 type of immune response over that of Th1. Effective stimulation of all the IgG isotypes, particularly IgG2a, after immunization with liposome encapsulated antigens seems to be responsible for the significant of resistance against the disease. Taken together, these data indicate a potential for the liposomal antigens as a vaccine which could trigger both humoral and cell-mediated immune responses.

58 THERE ARE 58 CITED REFERENCES AVAILABLE FOR THIS REFERENCE COUNT: RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L12 ANSWER 12 OF 16 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 1995:254536 CAPLUS DOCUMENT NUMBER: 122:38664

ORIGINAL REFERENCE NO.: 122:7363a,7366a

TITLE: Interactions in model vaccines composed of mixtures of

aluminum-containing adjuvants AUTHOR(S):

Al-Shakhshir, Ragheb H.; Lee, Ann L.; White, Joe L.; Hem, Stanley L.

CORPORATE SOURCE: Dep. Industrial and Physical Pharmacy, Purdue Univ.,

West Lafayette, IN, 47907, USA Journal of Colloid and Interface Science (1995), SOURCE:

169(1), 197-203

CODEN: JCISA5; ISSN: 0021-9797 Academic

PUBLISHER: DOCUMENT TYPE: Journal LANGUAGE: English

AB The optimum formulation of vaccines containing multivalent antigens may require that more than a single type of aluminum-containing adjuvant be used. In some cases, in order to maximize the binding of the neg. charged antigen(s), a pos. charged

adjuvant such as aluminum hydroxide could be used. In other

cases, if the antigen(s) were pos. charged,

a neg. charged adjuvant such as aluminum phosphate might be preferred. The multivalent vaccine would therefore be prepared by combining the individual monovalent bulks resulting in a suspension consisting of mixed aluminum-containing adjuvants. Studies of such mixed suspensions revealed that some phosphate ions from the aluminum phosphate

adjuvant desorbed upon the dilution which occurred when the

monovalent bulks were combined. The desorption of phosphate reduced the neg. surface charge of the aluminum phosphate adjuvant. The desorbed phosphate anions were subsequently readsorbed by the aluminum hydroxide adjuvant resulting in a decrease of its pos. surface charge. Description of the adsorbed antigens may also occur when the monovalent suspensions are mixed. In the model system studied, a significant fraction (25%) of adsorbed lysozyme desorbed from the aluminum phosphate adjuvant upon dilution (1:2). In contrast, almost no bovine serum albumin was desorbed from an aluminum hydroxide adjuvant upon similar dilution. A method based on measuring the electrophoretic mobility of the adjuvants was developed to assess the interactions that take place between the different adjuvants. Rapid aggregation was observed for the system consisting of oppositely charged adjuvants. The rate of aggregation of the pos. charged aluminum hydroxide adjuvant with the neg. charged aluminum phosphate adjuvant was reduced by the adsorption of proteins. Colloidal stability was enhanced by increased surface coverage of the proteins on the adjuvants. It was concluded that protein adsorption reduces the rate of aggregation of the mixed adjuvant system by minimizing the difference in surface charge between the aluminum-containing adjuvants and by providing steric repulsion.

L12 ANSWER 13 OF 16 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 1991:478692 CAPLUS DOCUMENT NUMBER: 115:78692

ORIGINAL REFERENCE NO.: 115:13435a,13438a

TITLE: The importance of surface charge in the optimization

of antigen-adjuvant interactions

AUTHOR(S): Callahan, Patricia M.; Shorter, Andrew L.; Hem,

Stanley L.

CORPORATE SOURCE: Dep. Pharm. Res., SmithKline Beecham Pharm., King of

Prussia, PA, 19406-0939, USA

Pharmaceutical Research (1991), 8(7), 851-8 SOURCE:

CODEN: PHREEB; ISSN: 0724-8741

DOCUMENT TYPE: Journal

LANGUAGE:

antigen-adjuvant interactions.

English The absorptive behavior of the recombinant malarial antigens R32et32, R32NS181, and NS181V20 to aluminum hydroxide and aluminum phosphate gels was studied as a function of pH and buffer ions. The Plasmodium falciparum antigen, R32NS181, and the P. vivax antigen , NS181V20, with isoelec. points (pI) of 5.9 and 5.5, resp., adsorbed readily to the pos. charged boehmite form of aluminum hydroxide gel. These two antigens displayed reversible, linear adsorption behavior in the pH range 5-9, with maximal adsorption observed at the lowest pH studied. The addition of acetate buffer ions had little effect on adsorption, while the presence of phosphate decreased adsorption for R32NS181 and NS181V20 by 24 and 40%, resp. The adsorptive behavior of these two antigens with the neg. charged adjuvant, aluminum phosphate, was markedly decreased. The converse situation was observed with the R32et32 antigen, whose pI is estimated to be 12.8. There was minimal interaction of this antigen with aluminum hydroxide gel except in the presence of phosphate counter ions and significant, nonreversible adsorption with aluminum phosphate gel. Enhanced adsorption of R32tet32 to aluminum hydroxide gel in the presence of phosphate is suggested to be the result of a covalent bond between a surface aluminum and a phosphate anion that modifies the surface charge of the aluminum hydroxide gel. These results indicate that the role of complementary surface charges, both for the ionization state of the protein and for the aluminum

adjuvants, is the key in optimizing conditions for significant

L12 ANSWER 14 OF 16 CAPLUS COPYRIGHT 2008 ACS on STN

1987:583417 CAPLUS ACCESSION NUMBER:

DOCUMENT NUMBER: 107:183417

ORIGINAL REFERENCE NO.: 107:29325a,29328a

TITLE: The effect of surface-coupled antigen of liposomes in

immunopotentiation

AUTHOR(S): Latif, Nahid Ali; Bachhawat, Bimal Kumar

Indian Inst. Chem. Biol., Calcutta, 700 032, India

SOURCE: Immunology Letters (1987), 15(1), 45-51

CODEN: IMLED6; ISSN: 0165-2478

DOCUMENT TYPE: Journal English

LANGUAGE:

The effect of surface coupled antigens of liposomes on the immunol.

response was investigated. Lysozyme was covalently coupled to neutral and pos. charged liposomes using glutaraldehyde. S.c. administration of these prepns. stimulated a significant antibody response higher than that elicited by the antigen entrapped in neutral liposomes. Immunization by liposomal antigens together with complete Freund's

adjuvant resulted in strong immune responses, highest with the

antigen coupled to neutral and pos. charged

liposomes followed by the antigen entrapped in neutral

liposomes. Primary and secondary immunization with lysozyme, both entrapped and coupled to liposomes, evoked an IgG response.

L12 ANSWER 15 OF 16 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 1987:520932 CAPLUS DOCUMENT NUMBER: 107:120932

ORIGINAL REFERENCE NO.: 107:19495a,19498a

Liposomes as immunological adjuvants:

antigen incorporation studies

Gregoriadis, Gregory; Davis, David; Davies, Alun AUTHOR(S): CORPORATE SOURCE: Sch. Med., R. Free Hosp., London, NW3 2QG, UK

SOURCE: Vaccine (1987), 5(2), 145-51 CODEN: VACCDE; ISSN: 0264-410X

DOCUMENT TYPE: Journal

LANGUAGE: English

Tetanus toxoid was incorporated into liposomes composed of equimolar phospholipid and cholesterol. The toxoid was either passively entrapped into multilamellar vesicles prepared by the dehydration-rehydration

procedure (DRV) or covalently coupled by diazotization to the surface of multilamellar vesicles (MLV) prepared by the classical procedure. Up to 82.3% of the antigen used was entrapped in neutral, neg. and

pos. charged DRV composed of a variety of unsatd. and

saturated phospholipids and 63.1% was coupled to MLV composed of egg phosphatidylcholine. After freeze-drying of toxoid-incorporating DRV and MLV and subsequent rehydration, up to 93.5% of the antigen was recovered with liposomes and, in the case of MLV, retained its external localization. Upon freeze-drying in the presence of 0.25M trehalose, up to 96.1% of the antigen was recovered with the DRV liposomes. In

immunization studies using Balb/c mice, DRV composed of equimolar egg phosphatidylcholine and cholesterol acted as immunol. adjuvants to the entrapped tetanus toxoid. In addition, there was no difference in

immune responses between DRV and MLV of identical composition but bearing the toxoid on their surface. A comparison of immune responses to the toxoid entrapped in DRV made of phospholipids with varying gel to liquid crystalline

transition temperature (Tc) revealed a reduction in responses to very low values for

DRV made of distearoylphosphatidylcholine (Tc 54°).

L12 ANSWER 16 OF 16 CAPLUS COPYRIGHT 2008 ACS on STN ACCESSION NUMBER: 1987:3570 CAPLUS

DOCUMENT NUMBER: 106:3570

ORIGINAL REFERENCE NO.: 106:695a,698a

TITLE: Effect of liposome charge on the intestinal absorption of macromolecules and on the induced immune response

Feknous, M.; Andre, F.; Andre, C. AUTHOR(S):

CORPORATE SOURCE: UER Biol. Hum., Univ. Claude Bernard, Lyon, Fr.

SOURCE: Medecine et Hygiene (1986), 44(1667), 2158, 2160-1,

2164-5 CODEN: MEHGAB; ISSN: 0025-6749

DOCUMENT TYPE: Journal

LANGUAGE: French

As a model of oral vaccination, human serum albumin was given to mice in the free form or encapsulated in multilamellar phosphatidylcholinecholesterol liposomes that were elec. neutral or charged neg. or pos. Compared with the intestinal absorption of the free albumin, that of the liposome-bound forms was increased 12-fold when the carrier was neutral and 3-fold when it was pos. or neg. charged. The immune response (as measured by the serum titer of antialbumin antibodies) was weak when the antigen was given in the free form or associated with the pos . charged liposome; it was greatly increased when the albumin

was associated with the neutral or, especially, the neg. charged liposome. This

enhancement of the immune response was probably due both to increased intestinal absorption and to the adjuvant effect of the neg. charged carrier.